

IN THE U.S. PATENT AND TRADEMARK OFFICE

Applicants: Eijiro WATANABE et al.

Serial No.: 08/992,914

Group: 1638

Filed: December 18, 1997

Examiner: D.H.Kruse

For: RAFFINOSE SYNTHASE GENES AND THEIR USE

DECLARATION UNDER 37 CFR 1.132

Honorable Commissioner of Patents and Trademarks
Washington, D.C. 20231

Sir:

I, Akitsu NAGASAWA, citizen of Japan and residing in Kamokogahara 3-28-56, Higashi-Nada-ku, Kobe-shi, Hyogo-ken, Japan, declare and say that:

1. I completed the master's course, with a major in agricultural biology, of the graduate school of Kyoto University and obtained a master's degree in agriculture at Kyoto University in March, 1984.

2. From April, 1984 to the present, I have been an employee of Sumitomo Chemical Company, Limited, the assignee of the above-identified application.

3. From April, 1984 to the present, I have been engaged in research works for plant engineering using recombination and other gene manipulation, such as cloning of plant genes, preparation and evaluation of transgenic plants.

4. I am one of the members of the research project related to the above-identified application and am familiar with the subject matter thereof.

5. I have read the Office Action mailed March 11, 2005 and the reference cited, and am familiar with the subject matter thereof.

6. To demonstrate successful identification of raffinose synthase genes in plant, I have made the following computer analysis.

ANALYSIS

1) The overall sequence homologies (%) among the amino acid sequences of raffinose synthases (RFSs), seed imbibition protein (SIP) and stachyose synthases (STSs) shown in Table 1 attached hereto were calculated based on a global multiple alignment (the alignment of sequences over their entire length) using the gene analysis software GENETYX-SV/RC for Windows version 6.1.0 (GENETYX Corporation; <http://www.sdc.co.jp/genetyx/>) with default parameters. The global multiple alignment was generated using CLUSTAL sequence analysis program. The amino acid sequences of the RFSs, SIP and STSs used to produce the global multiple alignment are as follows:

Sc-02:

MAPPSITKTATLQDVISTIDIGNGNPLFSITLDQSRDFLANGHPFLTQV
PPNITTNTTASSFLNLSNKDTIPNNNNTMLLQQGCFVGFNSTEPKSH
HVVPLGKLKGIKFMSIFRFKVWWTTHWVGTVNGQELQHETQMLILDKNDSL
GRPYVLLLPILENTFRDSLQPGLNDHIGMSVESGSTHVTGSSFKACLYIH
LSNDPYSILKEAVKVIQTQLGTFKTLEEKTAPSIIDKFGWCTWDAYLKV
HPKGVWEGVKSLTDDGCPPGFVIDDGWQSICHDDDEDDSGMNRTSAGE
QMPCRLVKYEENSFKFREYENPENGGKKGLGGFVRDLKEEFGSVESVYWH
ALCGYWGGVRPGVHGMPKARVVVPKVSQGLKMTMEDLAVDKIVENGVGLV
PPDFAHEMFDFGLHSHLESAGIDGVKVDVIHLLELLSEYGRVELARAYY
KALTSSVKKHFKGNGVIASMEHCNDFFLLGTEAISLGRVGDDFWCSDPG
DPNGTYWLQGCHMVHCAYNSLWMGNFIQPDWDMFQSTHPCAEFHAASRAI
SGGPIVYSDCVGNHNFKLLKSLVLPDGSILRCQHYALPTRDCLFEDPLHN
GKTMLKIWNLNKYTGVLGLFNCQGGGWCPEARRNKSVSEFSRAVTCYASP
EDIEWCNGKTPMSTKGVDFFAVYFFKEKKLRLMKCSDRLKVSLEPFSFEL
MTVSPVKVFSKRFIQFAPIGLVNMLNSGGAIQSLEFDDNASLVKIGVRGC
GEMSVFASEKPVCCKIDGVVKVFLYEDKMARVQILWPSSTLSLVQQLF

Sc-03:

MAPSFSKENSKTCDDEVANHDDCNTCPISLEESNFMVNNGHVILSQVPSNI
TAISKMGFDGLFVGFDAPEPKARHVSVGQLKGIPFMSIFRFKVWWTTHW
TGSNGRDLEHETQILILDKSDEGLGRPYIVILPLIEGPFRASLQPGSVDD
YVDICVESGSTKVGDSFRAVLYIRAGPDFKLIKDTMKEVQAHLGTFKL
LDDKTPPGIVDKFGWCTWDAYLKVEXYGVWEGVKGLENQVPPGLVILD
DGWQSICHDDDPITDQEGINRTSAGEQMPCRLIKYEENFKFRDYKSPNIM
GHEDHPNMGMRAFVRDLKEEFKTVEHVVYVHAFTGYWGGVRPNVPGLEXA
QVVTPLSPGLEMTMEDLAVDKIVNNGIGLVQPDKAQELYEGLHSHLENC

GIDGVKVDV IHLLEMMAEDYGGRVELAKTYYKAI TESVRKHFKGNGVI AS
MEQCNDMFMLLGTE I CLGRVGDDFWPTDPSGDINGTYWLQGCHMVHCAYN
SLWMGNF I HPDWDMFQSTHPCAEFHAASRAISGGPIYVSDVVGKHNIPLL
KRLVLADGSILRCEYHALPTKDCFLVDPLHDGKMLKIWNLNKYNGVLGV
FNCQGGGWSRESRKNLCFSEYSKPISCKTSPKDVEWENGHKPFP IKGVEC
FAMYFTKEKKILSQLSDTIEISLDPFDYELIVVSPMTILPWESIAFAPI
GLVNMLNAGGAVKSLDISEDNEDKMQVQVGIKGAGEMMVYSSSEKPKACRVN
GEDMEFEYEESMIKVQVTWNHNSGGFTTVEYLF

Sc-04 (truncated) :

MAPSISKTVELNSFGLVNGNLPLSITLEGSNFLANGHPLTEVPENIIVT
PSPIDAKSSKNNEDDVVGCFVGFADEPRSRHVASLGKLRGIKFMSIFR
FKVWWTTTHWVGSNGHELEHETQMMLLDKNDQLGRPVLILPILQASFRAS
LQPGDDYVDVCMESGSTRVCSSFGSCLYVHVCHDPYQLLREATKVRM
HLGTFKLLEEKTAPEVIIDKFGWCTWDAYLKVHPSGVWEGVKGLVEGGCP
PGMVLIDDGWQAI CHDEDPITDQEGMKRTSAGEQMPCCRVLKLEENYKFRQ
YCSGKDSEKGGMGAFVRDLKEQFRSVEQVYVWHALCGYWGGVRPKVPGMPQ
AKVVTPLSNGLKLTMKDLAVDKIVSNGVGLVPPHLAHLLYEGLHSRLES
AGIDGVKVDV IHLLEMSEEYGGRVELAKAYYKALTASVKKHFKGNGVI A
SMEHCNDFLLGTEAI ALGRVGDDFWCTDPSGDPNGTYWLQGCHMVHCAY
NSLWMGNF I QPDWDMFQSTHPCAEFHAPLGPSLVDQFTLVIVLESTSSC
SRASLCLMGRFCVVNTMHSPHETVCLKTPCMMGRQCSKFGISTNIQVFWV
YLIAKEVGGVP

Sc-05 :

MAPPSVIKSDAAVNGIDL SGKPLFRLEGS DLLANGHVLT DVPVNVTVA
SPYLA DKDGE PV DASAGSF IGFNLDGE PRSRHV ASI GKL RDIFM SIFR
FKVWWTTTHWVGSKGS DIENETQI I ILENSGSGRPYVLLLPLLEGSFRSSFQ
PGEDDDAVCVESGSTQVTGSEFRQVYVHAGDDPFKLVKDAMKVVRVHM
NTFKLLEEKXPPGIVDKFGWCTWDAYLTVNPDGVHKGVKCLVDGGCPPG
LVLIDDGWQSIGHDSGIDVEGMSCTVAGEQMPCCRLLKFQENFKFRDYVS
PKDKNEVGMKA FVRDLKEEFSTVDYIYVWHALCGYWGLRPGAPTLPPST
IVRPELSPGLKLTMQDLAVDKIVDTIGFVSPDMANEFYEGLHSHLQNVG
IDGVKVDV IHLLEMCEKYGGRVDLAKAYFKALTSSVNKHFDGNGVIASM
EHCNDFMFLGTEAI SLGRVGDDFWCTDPSGDI NGTYWLQGCHMVHCAYNS
LWMGNF I QPDWDMFQSTHPCAEFHAASRAISGGPIYISDCVGQHDFDLLK
RLVLPDGSILRCEHYALPTRDRLFEDPLHDGKMLKIWNLNKYTGII GAF
NCQGGGWC RETRRNQCF SQCVNTLTATTNPKDVEWNSGNNPISVENVEEF
ALFLSQSKKLVLSGPNDDLEITLEPFKFELITVSPVVTIEGSSVQFAPIG
LVNMLNTSGAIRSLVYHEESVEIGVRGAGEFRVYASRKPASCKIDGEVVE
FGYEE SVMVQVPWSAPEGLSSIKYEF

PsRFS :

MAPPSITKTATQQDVISTVDIGNSPLLSISLDQSRNFLVNGHPFLTQVPP
NITTTTSTPSPLDFKSNKDTIANNNTLQQQGCFVGFTNTEAKSHVV
PLGKLKGIKFTSIFRFKVWWTTHWVGTNQHELQHETQILILDKNISLGRP
YVLLPILENSFRSLQPGLNDYVDMVESGTHVTGSTFKACLYLHLSN
DPYRLVKEAVKVIQTKLGTFTLEEKTPPSIEKFGWCTWDAFYLKVKHPK
GVWEGVKALTDDGCPPGFVIDDGWQSISHDDDPVTERDGMNRTSAGEQ
MPCRLIKYEENYKFREYENGDNGGKGLVGFVRDLKEEFRSVEVYVWHA
LCGYWGGRPKVCGMPEAKVVVPKLSPGVKMTMEDLAVDKIVENGVGLVP
PNLAQEMFDGIHSHLESAGIDGVKVDVIHLLELLSEYGGRVELAKAYYK
ALTSSVNKHFKGNGVIASMEHCNDFFLLGTEAISLGRVGDDFWCCDPSGD
PNGTYWLQGCHMVHCAYNSLWMGNFIHPDWDMFQSTHPCAEFHAASRAIS
GGPVYVSDCVGNHNFKLLKSFVLPDGSLRQCQHYALPTRDCLFEDPLHNG
KTMKCIWNLNKYAGVGLFNCQGGGWCETRRNKSASEFSHAVTCYASPE
DIEWCNGKTPMDIKGVDVFAVYFFKEKKLSLMKCSRDRLEVSLEPFSFELM
TVSPLKVF SKRLIQFAPICLVNMLNSGGAVQSLEFDDASLVKIGVRGCG
ELSVFASEKPVCCKIDGVSVF DYEDKMVRVQILWPGSSTLSLVEFLF

Aj-05 :

MAPSFKNNGGSNVVSDGLNDMSSPFAIDGSDFTVNGHSFLSDVPENIVAS
PSPYTSIDKSPVSVGCFVGF DASEPDSRHVVSI GKLKDIFRMSIFRFKVW
WTTHWVGRNGGDLESETQIVILEKSDSGRPYVFLPIVEGPFRTSIQPGD
DDFVDVCVESGSSKVVDASFRSMLYLHAGDDPFALVKEAMKIVRTHLGT
RLLEEKTPPGIVDKFGWCTWDAFYLTVHPQGVIEGVRHLVDGGCPPGLVL
IDDGWQSIGHDSDPITKEGMNQTVAGEQMP CRLKQFQENYKFRDYVNPKA
TGPRAGQKGMKAFIDELKGEFKTVEHVYVWHALCGYWGLRPQVPGLEA
RVIQPVLSPLQLQMTMEDLAVDKIVLHKVGLVPPEKAEMYEGLHAHLEKV
GIDGVKIDVIHLLEMLCEDYGGRVDLAKAYYKAMTKSINKHFKGNGVIAS
MEHCNDMFLGTEAISLGRVGDDFWCTDPSGDPNGTFWLQGCHMVHCAND
SLWMGNFIHPDWDMFQSTHPCAAFHAASRAISGGPIYVSDSVGKHNF DLL
KKLVLPDGSLRSEYYALPTRDCLFEDPLHNGETMLKCIWNLNKFTGVIGA
FNCQGGGWCETRRNQCSQYSKRVTSKTNPKDIEWHSGENPISIEGVKT
FALYLYQAKKILSKPSQDLDIALDPFEFELITVSPVTKLIQTSLHFAP
GLVNMLNTSGAIQSVDYDDDLSSVEIGVKGC GEMRVFASKPRACR IDGE
DVGFKYDQDQM VVQWPWIDSSGGISVIEYLF

HvSIP :

MTVTPQITVGDGRLAVRGRTVLSGVPDNVTAHAAGAGLVDGAFVGATAA
EAKSHHVFTFGTLRDCRFMCLFRKLWWMTQRMGTSGRDVPLETQFILIE

VPAAAGNDDGDSSDGDSEPVYLVMLPLEGQFRVLQGNDQDELQICIES
GDKAVETEQGMNNVYVHAGTNPFDTITQAVKAVEKHTQTFHHREKKTVP
FVDWFGWCTWDAYTDVTADGVKQGLRSLAEGGAPRFLIIDDGWQQIGS
ENKDDPGVAVQEGAQFASRLTGIRENTKFQSEHNQEETPGLKRLVDET
EHGVKSYYVWHAMAGYGGVKGPSAAGMEHYEPALAYPVQSPGVTGNQPD
VMDSLSVLGLGLVHPRRVHRYDELHAYLAACGVGVKVDVQNI
VETLGA
GHGGRVALTRAYHRALEASVARNFPDNGCISCMCHNTDMLYSAKQTAV
ASDDFYPRDPASHTVHISSVAYNTLFLGEFMQPDWDMFHSLHPAAEYHGA
ARAIGGCPIVYSDKPGNHNFLLRKVLVDGSVLRQLPGRPTRDCLFSD
PARDGASLLKIWNMNKCAVVGVFCNCQGAGWCRVAKKTRIHDEAPGTLTG
SVRAEDVEAIQAAGTGDWGGEAVVYAHRAGELVRLPRGATLPVTLKRE
YELFHVCVPVRAVAGVSFAPIGLLHMFNAGGAAVEECTVETGEDGNAVV
RVRGCGRGAYCSRRPAKCSVDSADVEFTYDSDTGLVTADVPVPEKEMY
CALEIRV

AmSTS :

MAPPYDPIPIPIPMSAILNFLSSTVKDNSFELLGTL SVKNPI LTDIPS
NVSFSSFSSIVQSSEAPVPLFQRAQSLSSSGFLGFSQNEPSSRLMNSLG
KFTDRDFVSIFRFKTTWSTQWVGGTTGSDIQMETQWIMLDVPEIKSYAVVV
PIVEGKFRSALFPKGDHILIGAESGTVKTSNFDAIAYVHSENPTYL
MRDAYTAVRVHLNTFKLIEEKSAPPLVNKFGWWTWDAFYLTVEPAGIYHG
VQEFADEGGLTPRFLIIDDGWQSINNDDNDPNEDAKNLVLLGGTQMTARLHR
LDECEKFRKYKGGSMSGPNRPFDPKKPKLLISKAIEIEVAEKARDAAQ
SGVTDLARYEAEIEKLTKELDQMFGGGGEETSSGKSCSSCSCKSDNFGMK
AFTKDLRTNFGLDDIYVWHALAGAWGGVVRPGATHNAKIVPTNLSPGLD
GTMTDLAVVKIEGSTGLVDPDQAEDFYDSMHSYLSVGITGVKVDVIHT
LEYISEDYGGRVELAKAYYKGLSKSLAKNFNGTGLISSMQQCNDFFLLGT
EQISMGRVGDDFWFQDPNGDPMGVYWLQGVHMIHCAYNSWMQFQI
DMFQSDHPGGYFHAGSRAICGGPVYVSDSLGGHNFLLKKLVFNDGTIPK
CIHFALPTRDCLFKNPLFDSKTIKIKWNFKYGGVIGAFNCQGAGWDPKE
QRIGKGSQCYKPLSGSVHVGSGIEFDQKKEASEMGEAEYYAVLSEAEL
LATRDSDPIKITIQSSTFEIFSFVPIKKLGEVKFAPIGLTNLFNAGGTI
QGLVYNEGIAKIEVKGDGKFLAYSSVVPKKAYVNGAEKVAWSGNGKLEL
DITWYEECGGISNVTFVY

PsSTS-1 :

MAPPLNSTTSNLIKTESIFDLSERKFKVKGFLFHDVPENVSFRSFSSIC
KPSESNAPPSSLQKVLAYSHKGFFGFSHETPSDRLMNSIGSFNGKDFLS
IFRFKTTWSTQWIGKGSIDLQMETQWILIEVPETKSYVVIIPIIEKCFRS
ALFPGFNDHVKIIAESGTVKESTFNSIAYVHSENPYDLMKEAYSAIR
VHLNSFRLLEEKTIIPNLVDKFGWCTWDAYLTVPNPIGIFHGLDDFSKGGV

EPRFV I IDDGWQS I SFDGYDPNEDAKNL VL GGEQMSGRL HRFDECYKFRK
YESGLLGPNSPYDPNNFTDL I LKG I EHEKLRKKREEA I SSKSSDLAE I
ESKIKKVVKE I DDLFGGEQFSSGEKSEMKE YGLKAFTKDLRTKFKGLDD
VYVWHALCGAWGGVRPETTHLDTK I VPCKLSPGLDGTMEDLAVVE I SKAS
LGLVHP SQANELYDSMHSYLAESG I TGVKVDV I HSLEYVCDEYGGRVDLA
KVYYEGLTKS I VKNFNGNGM I ASMQHCNDFFLGTKQ I SMGRVGDDFWFQ
DPNGDPMGSFWLQGVHM I HCSYNSLWMQGM I QPDWDMFQSDHVCASFHAG
SRAICGGP I YVSDNVGSHDFDL I KKLVFPDGT I PKC I YFPLPTRDCLFKN
PLFDHTTVLK I WNFNKG YGGV I GAFNCQGAGWDP I MQKFRGFPECYKP I PG
TVHVTVEWDQKEETSHLGKAEEYVYLNQAEELSLMTLKSEP I QFT I QP
STFELYSFVPTKLCGG I KFAP I GLTNMFNSGGTVIDLEYVGNGAK I KV
GGGSFLAYSSES PKKFQLNGCEVDFEWLGDGKLCVNVPWIEEACGVSDME
IFF

PsSTS-2:

MAPPLNSTTSNL I KTES I FDL SERKFKVKGFLFHDV PENV SFRS FSS I C
KPSESNAPPSSLQKVLA YSHKGFFGF SHETPSDRLMNSLGSFNGKDFLS
I FRFKTWWSTQW I GKSGSDLQMETQW I LIEVPETKSYVV I I P I I EKCFRS
ALFPGFNDHV K I I AEGSTKVKESTFNS I AYVHFSENPYDLMKEAY I A I R
VHLNSFRLLEEKT I PNLVDKFGWCTWDAFYLTVNP I G I FHGLDDFSKGGV
EPRFV I IDDGWQS I SFDGCDPNEDAKNL VL GGEQMSGRL HRFDECYKFRK
YESGLLGPNSPYDPKKFTDL I LKG I EHEKLRKKREEA I SSKSSDLAE I
ESKIKKVVKE I DDLFGGEQFSSVEKSEMKE YGLKAFTKDLRTKFKGLDD
VYVWHALCGAWGGVRPETTHLDTKFVPCKLSPGLDGTMEDLAVVE I SKAS
LGLVHP SQANELYDSMHSYLAESG I TGVKVDV I HSLEYVCDEYGGRVDLA
KVYYEGLTKS I VKNFNGNGM I ASMQQCNDFFLGTKQ I SMGRVGDDFWFQ
DPNGDPMGSFWLQGVHM I HCSYNSLWMQGM I QPDWDMFKSDHVCASFHAG
SRAICGGP I YVSDNVGSHDFDL I KKLVFPDGT I PKC I YFPLPTRDCLFKN
PLFDHTLLK I WNFNKG YGGV I GAFNCQGAGWDP I MQKFRGFPECYKP I PG
TVHVTQVEWDQKEETSHFGKAEEYVYLNQAEELCLMTLKSEP I QFT I QP
STFELYSFVPTKLCGG I KFAP I GLTNMFNSGGTVIDLEYVGNGAK I KV
GGGSFLAYSSES PKKFQLNGCEVDFEWLGDGKLCVNVPWIEEACGV

SaSTS:

MAPPNDP I SS I FSPL I SVKKDNAFEL VGGKLSVKNVPLLSE I PSNVTFKS
FSS I CQSSGAPAPLYNRAQSLSCGGFLGSQKESADSVTNSLGKFTNRE
FVS I FRFKTWWSTQWVGTSGSD I QMETQW I MLNLP I KSYAVV I I VEGK
FRSALFPGKDGHVL I SAESGSTCVKTTSFTS I AYVHVS DNPYTLMDGYT
AVRVHLDTFKL I EEKSAAPLVNKFGWCTWDAFYLTVEPAG I WNGVKEFSD
GGFSRFL I IDDGWQS I NIDGQDPNEDAKNL VL GGTQMTARL HRFDECEK
FRKYKGSMGPKVYFDPKKPKLL I SKA I E I EGVEKARDKA I QSG I TDL

SQYE IKLKKLNKELEMFGGGNDEKGSSKGSCDCSCKSQNSGMKAFTND
LRTNFGLDDIYVWHALAGAWGGVKPGATHLNAKIEPCKLSPGLDGTMD
LAVKILEGSIGLVHPDQAEDFYDSMHSYLSKVGITGVKVDVIHTLEYVS
ENYGGRVELGKAYYKGLSKSLKKNFNGSGLISSMQQCNDFFLLGTEQISM
GRVGDDFWFQDPNGDPMGVFWLQGVHMIHCAYNSMWMGQIHPDWDMFQS
DHCSAKFHAGSRAICGGPVYVSDSLGGHDFDLLKKLVFNDGTIPKCIHFA
LPTRDCLFKNPLFDSKTIKIKWNFNKYGGVVGAFNCQGAGWDPKEQRICKG
YSECYKPLSGSVHVSDEWDQKVEATKMGAEEYAVYLTSEKLLTTPE
SDPIPFTLKSTTFEIFSFVPIKKLGQGVKFAPIGLTNLFNSGGTIQGVVY
DEGVAKIEVKGDGKFLAYSSVPKRSYLNGEVEYKWSGNKGVEVDVPWY
EECGGISNITFVF

VaSTS:

MAPPNDPVNATLGLEPSEKVFDSLGDKLTVKGVVLLSHVPENVTSSFSS
ICVPRDAPSSILQRVTAASHKGGLGFSHVSPSDRLINSLGSFRGRNFLS
IFRFKTTWSTQWVGNNSGSDLQMETQWILIEVPETESYVVIIPIIEKSFRS
ALHPGSDDHVKICAESGSTQVRASSFGIAAYVHVAETPYNLMREAYSALR
VHLDLSFRLLEEKTVPRIVDKFGWCTWDAYLTVNPGVWHLKDFSEGGV
APRFVVIDDGWQSVDNFDDEDPNEDAKNLVLGGEQMTARLHRFEEGDKFRK
YQKGLLLGPNAPSFPETIKELISKGIEAEHLGKQAAIASAGGSDLAEIE
LMIVKVREEIDDLFGGKGKESNESGGCCCKAAECGGMKDFTTDLRTEFKG
LDDVYVWHALCGGWGGVRPGTTHLDSKIPCKLSPGLVGTMKDLAVDKIV
EGSIGLVLPHQANDLYDSMHSYLAQTVGTGVKIDVIHSLEYVCEEYGGRV
EIAKAYDGLTNSIKNFNGSGIIASMQQCNDFFLGTKQIPFGRVGDDF
WFQDPNGDPMGVFWLQGVHMIHCSYNSLWMGQIQPDWDMFQSDHECAKF
HAGSRAICGGPVYVSDSVGSHDFDLIKKLVFPDGTVPKCIYFPLPTRDCL
FRNPLFDQKTVLKIKWNFNKYGGVIGAFNCQGAGWDPKGKKFGFPECYKA
ISCTVHVTVEWDQKKEAEHMGKAEEYVYVLNQAEVLHLMTPVSEPLQLT
IQPSTFELYNFVPEKLGSSNIKFAPIGLTNMFNSGGTIQELEYIEKDVK
VKVKGGGRFLAYSTQSPKKFQLNGSDAAFQWLPDGKLTNLAWIEENDGV
SDLAIFF

The calculated overall sequence homologies (%) are shown in Table 2 attached hereto. The homologies between RFSs and SIP are less than 40%. The homologies between RFSs and STSs are not higher than 45%. On the other hand, the homologies among RFSs are all 50% or higher. Thus, the homologies among RFSs are higher than those homologies between RFSs and SIP and between RFSs and STSs.

A molecular phylogenetic tree of the RFSs, SIP and STSs shown in Table 1 is

drawn in Figure 1 attached hereto. The molecular phylogenetic tree is drawn by the UPGMA method using the gene analysis software GENETYX-SV/RC for Windows version 6.1.0 (GENETYX Corporation; <http://www.sdc.co.jp/genetyx/>) with default parameters. In the molecular phylogenetic tree, RFSs, SIP and STSs form different groups respectively.

In summary, Table 2 and Figure 1 show that RFSs, SIP and STSs can be distinguished from one another based upon a comparison of their amino acid sequences.

2) Attached Table 3 shows the identities obtained using the BLAST program for the amino acid sequences of RFSs, SIP and STSs shown in Table 1. Among Sc-02, Sc-03, Sc-04 and Sc-05, the identities were obtained by searching the "patent database" provided by NCBI (National Center for Biotechnology Information) with default parameters, using the amino acid sequence of each protein as the "query", and using "Protein query vs. translated database (tblastn)" of the NCBI BLAST program. Also, other identities were obtained by searching the "non-redundant database" provided by NCBI with default parameters, using the amino acid sequence of each protein as the "query", and using "Protein-protein BLAST (blastp)" of the NCBI BLAST program. The above-identified amino acid sequences of the RFSs, SIP and STSs are used as the "query" except that the amino acid sequence of Sc-04 used as the "query" is as follows:

Sc-04 (full-length) :

MAPSISKTVELNSFGLVNGNLPLSITLEGSNFLANGHPFLTEVPENIIVT
PSPIDAKSSKNNEDDVVGCFVGFHADEPRSRHVASLGKLRGIKFMSIFR
FKVWWTTHWVGNSNGHELEHETQMMLLDKNDQLGRPFVILPILQASFRAS
LQPGLDDYVDVCMESGSTRVCGSSFGSCLYVHVGHDPYQLLREATKVVRM
HLGTFKLLEEKTAPEVIIDKFGWCTWDAYLKVPHPSGVWEGVKGLVEGGCP
PGMVLIDDGWQAICHDEDPTIDQEGMKRTSAGEQMPCRLVKLEENYKFRQ
YCSGKDSEKGMGAFVRDLKEQFRSVEQVYVWHALCGYWGGVRPKVPGMPQ
AKVVTPKLSNGLKLTMKDLAVDKIVSNGVGLVPPHLAHLLYEGLHSRLES
AGIDGVKVDVIHLLEMLSEEVGGRVELAKAYYKALTASVKKHFKGNGVIA
SMEHCNDFFLLGTEIALGRVGDDFWCTDPGDPNGTYWLQGCHMVHCAY
NSLWMGNFIQPDWDMFQSTHPCAEFHAASRAISGGPVYVSDCVGKHNFKL
LKSLALPDGTILRCQHYALPTRDCLFEDPLHDGKMLKIWNLNKYTGVLG

LFNCQGGGWCPVTRRNKSASEFSQTVTCLASPQDIEWNGKSPICIKGMN
VFAVYLFKDHLKLMKASEKLEVSLEPFTFELLTVSPVIVLSKKLIQFAP
IGLVNMLNTGGAIQSMEFDNHIDVVKIGVRGCGEMKVFASEKPVSCLDG
VVVKFDYEDKMLRVQVPWPSASKLSMVEFLF

As shown in Table 3, the identities between RFSs and SIPs are about 40%. The identities between RFSs and STSs range from about 40% to about 50%. On the other hand the identities among RFSs are 60% or higher. The identities among STSs are also 60% or higher. That is, the identities among RFSs or the identities among STSs are higher than the identities between RFSs and SIP or the identities between RFSs and STSs. Thus, RFSs, SIP or STSs can be distinguished based on the results of analysis using BLAST program.

3) Attached Table 4 shows the identities obtained using another BLAST program for the amino acid sequences of RFSs, SIP and STSs shown in Table 1. All possible pair-wised amino acid sequence comparison were made by the "Blast 2 Sequences" program from NCBI (<http://www.ncbi.nlm.nih.gov/blast/bl2seq/bl2.html>). Sequence identities were calculated using default parameters, program; blastp, matrix; BLOSUM62, open gap penalty; 11, extension gap penalty; 1, gap x_dropoff; 50, expect; 10.0, and word size; 3. The amino acid sequences of the RFSs, SIP and STSs used to calculate sequence identities are identical to those used as the "query" to obtain identities shown in Table 3. Results were essentially the same with former two types of comparison.

4) In conclusion, raffinose synthases (RFSs), seed imbibition protein (SIP) and stachyose synthases (STSs) were clearly distinguished from one another based on comparison of their amino acid sequences.

7. I declare further that all statements made herein of my own knowledge are true and that all statements made on information and belief are believed to be true; and further that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonments, or both, under Section 1001 of Title 18 of the United States Code and that such willful false statements may jeopardize the validity of the above-identified application or any patent issued thereon.

This 6th day of September, 2005



Akitsu NAGASAWA

Table 1

Code	Protein*	Organism	Accession**	Reference	Author/Assignee
Sc-03	RFS	<i>Beta vulgaris</i>	E37133	09/301,766	Sumitomo Chemical
Sc-05	RFS	<i>Brassica juncea</i>	E36417	09/301,766	Sumitomo Chemical
Sc-02	RFS	<i>Vicia faba</i>	E24423	08/992,914	Sumitomo Chemical
Sc-04	RFS	<i>Glycine max</i>	E24424	08/992,914	Sumitomo Chemical
Aj-05	RFS	<i>Cucumis sativus</i>	AF073744	Family GH36***	Ohsumi et al.
PsRFS	RFS	<i>Pisum sativum</i>	AJ426475	Family GH36	Peterbauer et al.
HvSIP	SIP	<i>Hordeum vulgare</i>	M77475	Family GH36	Heck et al.
PsSTS-1	STS	<i>Pisum sativum</i>	AJ311087	Family GH36	Peterbauer et al.
PsSTS-2	STS	<i>Pisum sativum</i>	AJ512932	Family GH36	Peterbauer et al.
VaSTS	STS	<i>Vigna angularis</i>	Y19024	Family GH36	Peterbauer et al.
AmSTS	STS	<i>Alonsoa meridionalis</i>	AJ487030	Family GH36	Voitsekhovskaja
SsSTS	STS	<i>Stachys affinis</i>	AJ344091	Family GH36	Pesch and Schmitz

*Protein: RFS, Raffinose synthase; SIP, Seed Imbibition Protein; STS, Stachyose synthase.

**Accession: GenBank Accession Number.

***Family GH36: glycoside hydrolase family 36 (see Carbohydrate-Active Enzymes (CAZy) database: http://afmb.cnrs-mrs.fr/CAZY/GH_36.html)

Table 2

Table 3

Table 4

Fig. 1

[GENETYX : Evolutionary tree]
Date : 2004.2.4
Method: UPGMA

